(FILE 'HOME' ENTERED AT 14:24:39 ON 26 APR 2005)

	FILE 'CAPL	JS' ENTERED AT 14:24:53 ON 26 APR 2005
L1	188	SEA ABB=ON PLU=ON ACETYLCHOLINE ESTERASE INHIBITOR
L2	53	SEA ABB=ON PLU=ON CHOLINE ESTERASE INHIBITOR
L3		SEA ABB=ON PLU=ON L1 OR L2
L4	48	SEA ABB=ON PLU=ON (TACRINE OR PHYSOSTIGMINE OR RIVMSTIGMINE
		OR GALANTNMINE OR CITICOLINE OR VELNACRINE MALEATE OR METRIFONA
		TE OR HEPTASTIGMINE) AND (L1 OR L2)
L5	0	SEA ABB=ON PLU=ON L4 AND (DRUG OR SUBSTANCE OR OPIOID OR
		OPIATE OR ALCOHOL OR MARIJUANA OR HEROINE OR PHENCYCLIDINE OR
		AMPHETAMINE OR COCAINE) (P) (ABUSE OR WITHDRAWAL OR DEPENDANCY
		OR DEPENDENCY)
L6	24	SEA ABB=ON PLU=ON L4 AND (DRUG OR SUBSTANCE OR OPIOID OR
		OPIATE OR ALCOHOL OR MARIJUANA OR HEROINE OR PHENCYCLIDINE OR
		AMPHETAMINE OR COCAINE)
		D LOG 1- IBIB KWIC
		DICIDID WIC 1

FILE HOME

L6 ANSWER 20 OF 24 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 1991

1991:223359 CAPLUS

DOCUMENT NUMBER:

114:223359

TITLE:

Involvement of the cholinergic neuronal system and

benzodiazepine receptors in alcohol-induced

amnesia

AUTHOR (S):

Nabeshima, Toshitaka; Tohyama, Keiko; Ishihara,

Seiichi; Kameyama, Tsutomu

CORPORATE SOURCE:

Fac. Pharm. Sci., Meijo Univ., Nagoya, 468, Japan

SOURCE:

European Journal of Pharmacology (1991), 195(2), 285-9

CODEN: EJPHAZ; ISSN: 0014-2999

DOCUMENT TYPE:

Journal

LANGUAGE:

English

TI Involvement of the cholinergic neuronal system and benzodiazepine receptors in alcohol-induced amnesia

The involvement of the GABAergic and cholinergic neuronal systems and benzodiazepine (BZP) receptors in ethanol-induced amnesia was investigated using a passive avoidance task. Pretraining administration of ethanol impaired the passive avoidance response. The BZP agonist chlordiazepoxide potentiated the amnesia, while the GABA antagonists bicuculline and picrotoxin failed to affect it. The acetylcholine esterase inhibitor physostigmine partially attenuated the ethanol-induced amnesia. These results suggest that ethanol-induced amnesia is related to BZP receptors and a dysfunction of the cholinergic neuronal system.

ST ethanol amnesia GABA benzodiazepine receptor; alc amnesia GABA benzodiazepine receptor; cholinergic neuron ethanol amnesia

IT 57-47-6, **Physostigmine** 9000-81-1, Acetylcholinesterase

RL: BIOL (Biological study)

(ethanol-induced amnesia response to)

CAPLUS COPYRIGHT 2005 ACS on STN ANSWER 20 OF 24 L6

ACCESSION NUMBER:

1991:223359 CAPLUS

DOCUMENT NUMBER:

114:223359

TITLE:

Involvement of the cholinergic neuronal system and

benzodiazepine receptors in alcohol-induced

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CORPORATE SOURCE:

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ethanol amnesia GABA benzodiazepine receptor; alc amnesia GABA ST benzodiazepine receptor; cholinergic neuron ethanol amnesia

9000-81-1, Acetylcholinesterase IT 57-47-6, Physostigmine

RL: BIOL (Biological study)

(ethanol-induced amnesia response to)

L7 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:314578 CAPLUS

DOCUMENT NUMBER: 132:318050

Choline esterase inhibitors, alone or with other

agents, for treating restless legs syndrome and/or periodic limb movements during sleep, and diagnostic

method

INVENTOR(S): Hedner, Jan; Kraiczi, Holger

PATENT ASSIGNEE(S): Swed.

SOURCE: PCT Int. Appl., 26 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

TITLE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE	
WO 2000025821	A1 2000	 0511 WO 1999-SE1979	19991103	
RW: AT, BE, CH,		ES, FI, FR, GB, GR, IE, IT		
PT, SE EP 1154795	A1 2001	1121 EP 1999-957453	19991103	

1154795 A1 20011121 EP 1999-957453 19991103 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

IE, FI PRIORITY APPLN. INFO.:

SE 1998-3760 A 19981104 WO 1999-SE1979 W 19991103

REFERENCE COUNT: 7 THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Amethod for treating or preventing the restless legs syndrome and/or the periodic limb movements during sleep comprises administration of a choline esterase inhibitor (CEI) and, optionally, carbamazepine, clonidine, baclofen, hypnotic agent, opioid agonist, and dopaminergic agonist. Administration precedes the onset of sleep at night by from zero to three hours so as to make the CEI exert a therapeutic effect during a major portion of the sleep period. Also disclosed are corresponding pharmaceutical compns. and their use, including compns. comprising a combination of CEI with carbamazepine, clonidine, baclofen, hypnotic agent, opioid agonist, and dopaminergic agonist.

IT. Antihistamines

Diagnosis

Dopamine agonists

GABA agonists

Hypnotics and Sedatives

Movement disorders

Test kits

(choline esterase inhibitors, alone or with other agents, for treating restless legs syndrome and/or periodic limb movements during sleep, and diagnostic method)

L13 ANSWER 13 OF 13 CAPLUS COPYRIGHT 2005 ACS on STN

1996:726271 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 126:26879 TITLE:

Neuronal nicotinic acetylcholine receptors in the

Vidal, Catherine; Changeux, Jean-Pierre AUTHOR (S):

Dept. Molecular Neurobiology, Institut Pasteur, Paris, CORPORATE SOURCE:

75724/15, Fr.

News in Physiological Sciences (1996), 11(Oct.), SOURCE:

202-208

CODEN: NEPSEY; ISSN: 0886-1714 American Physiological Society

PUBLISHER: DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review, with 15 refs. Recent mol., immunol., and physiol. studies have AΒ revealed that a wide variety of nicotinic acetylcholine receptors exist in the nervous system of vertebrates. Nicotinic systems

in the brain appear to play significant roles in drug

addiction and in cognitive functions, as well as in pathologies,

such as Alzheimer's disease.

L13 ANSWER 11 OF 13 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:595513 CAPLUS

DOCUMENT NUMBER: 133:306385

Genetic dissection of nicotine-related behavior: a TITLE:

review of animal studies

AUTHOR (S): Mohammed, Abdul H.

CORPORATE SOURCE: Division of Geriatric Medicine, NEUROTEC, Karolinska

Institutet, Huddinge University Hospital, Huddinge,

S-141 86, Swed.

Behavioural Brain Research (2000), 113(1,2), 35-41 SOURCE:

CODEN: BBREDI; ISSN: 0166-4328

Elsevier Science Ireland Ltd. PUBLISHER:

Journal; General Review DOCUMENT TYPE:

English LANGUAGE:

THERE ARE 70 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 70

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

A review and discussion with 70 refs. Nicotine has a broad spectrum of behavioral effects. A considerable body of data has emerged indicating genetic factors regulate the behavioral effects of nicotine. Exptl. genetic techniques have been invaluable in generating knowledge on the interrelationship of genetic factors and behavioral responsiveness to nicotine. Three different approaches have been invoked to explore the relationship of genetic factors in response to nicotine. Firstly, the classical genetic tool of inbred lines has been exploited to delineate genetic influences in the effects of nicotine. Secondly, the use of selectively bred lines has been profitably employed to reveal genetic differences in behavioral responses, such as cognition and exploration, to nicotine. These approaches have also provided useful information on the contribution of genetic factors influencing nicotinic receptors function. Finally, the mol. genetic technique of gene targeting to create mice with null mutations of specific genes in the central nervous system, which is having a tremendous impact in drug addiction research, has also been employed to gain insight into the mol. and cellular basis of nicotine action. These techniques are proving to be invaluable in dissecting the role of different subunits of the nicotinic acetylcholine receptors on behavior. This paper provides a survey of the animal studies that have used the above mentioned techniques to gain insight into the genetic basis of the behavioral effects of nicotine.

L13 ANSWER 10 OF 13 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2000:663192 CAPLUS

DOCUMENT NUMBER: 134:157453

TITLE: The development and expression of locomotor

sensitization to nicotine in the presence of ibogaine

AUTHOR(S): Zubaran, C.; Shoaib, M.; Stolerman, I. P.

CORPORATE SOURCE: Section of Behavioural Pharmacology, King's College

London, London, UK

SOURCE: Behavioural Pharmacology (2000), 11(5), 431-436

CODEN: BPHAEL; ISSN: 0955-8810

PUBLISHER: Lippincott Williams & Wilkins

DOCUMENT TYPE: Journal LANGUAGE: English

REFERENCE COUNT: 32 THERE ARE 32 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

Ibogaine is a naturally occurring psychoactive alkaloid with claimed efficacy in the treatment of certain drug addictions, including nicotine. It was reported to be a non-competitive blocker of nicotinic receptors, with a potent inhibitory action on nicotinic acetylcholine receptor-mediated catecholamine release. The authors have investigated the effect of different doses of iboqaine on the development and expression of sensitization to the locomotor stimulant effect of nicotine in rats, a facilitatory process in which a history of exposure to nicotine results in enhanced locomotor activity when the same dose of nicotine is administered repeatedly. The effects were determined of co-administering ibogaine (0.0, 5.0 or 10 mg/kg i.p.) with nicotine (0.0 $\,$ or 0.4 mg/kg s.c.) daily for 21 days. Dose-response curves for nicotine (0.04-0.8 mg/kg s.c.) were then determined in groups of 10 rats. There was clear sensitization of the locomotor activity produced by nicotine in photocell activity cages but co-administration of ibogaine with nicotine had no effect on the degree of sensitization. Ibogaine (5-20 mg/kg) itself did not influence locomotor activity and was also without effect on the expression of the sensitized response to 0.4 mg/kg of nicotine (n = 10). Thus, there was no evidence that iboqaine may retard or suppress sensitization to nicotine

L13 ANSWER 7 OF 13 CAPLUS COPYRIGHT 2005 ACS on STN

ACCESSION NUMBER: 2002:641500 CAPLUS

DOCUMENT NUMBER: 137:278435

TITLE: Evidence that intermittent, excessive sugar intake

causes endogenous opioid dependence

AUTHOR(S): Colantuoni, Carlo; Rada, Pedro; McCarthy, Joseph;

Patten, Caroline; Avena, Nicole M.; Chadeayne, Andrew;

Hoebel, Bartley G.

CORPORATE SOURCE: Department of Psychology, Princeton University,

Princeton, NJ, USA

Obesity Research (2002), 10(6), 478-488

CODEN: OBREFR; ISSN: 1071-7323

PUBLISHER: North American Association for the Study of Obesity

DOCUMENT TYPE: Journal LANGUAGE: English

sugar-dependent

SOURCE:

REFERENCE COUNT: 73 THERE ARE 73 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT Objective: The goal was to determine whether withdrawal from sugar can cause AB signs of opioid dependence. Because palatable food stimulates neural systems that are implicated in drug addiction, it was hypothesized that intermittent, excessive sugar intake might create dependency, as indicated by withdrawal signs. Research Methods and Procedures: Male rats were food-deprived for 12 h daily, including 4 h in the early dark, and then offered highly palatable 25% glucose in addition to chow for the next 12 h. Withdrawal was induced by naloxone or food deprivation. Withdrawal signs were measured by observation, ultrasonic recordings, elevated plus maze tests, and in vivo microdialysis. Results: Naloxone (20 mg/kg i.p.) caused somatic signs, such as teeth chattering, forepaw tremor, and head shakes. Food deprivation for 24 h caused spontaneous withdrawal signs, such as teeth chattering. Naloxone (3 mg/kg s.c.) caused reduced time on the exposed arm of an elevated plus maze, where again significant teeth chattering was recorded. The plus maze anxiety effect was replicated with four control groups for comparison. Accumbens microdialysis revealed that naloxone (10 and 20 mg/kg i.p.) decreased extracellular dopamine (DA), while dose-dependently increasing acetylcholine (ACh). The naloxone-induced DA/ACh imbalance was replicated with 10% sucrose and 3 mg/kg naloxone s.c. Discussion: Repeated, excessive intake of sugar created a state in which an opioid antagonist caused behavioral and neurochem. signs of opioid withdrawal. The indexes of anxiety and DA/ACh imbalance were qual. similar to withdrawal from morphine or nicotine, suggesting that the rats had become

Searches for User *lchannavajjala* (Count = 10260)

Queries 10211 through 10260.

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S#	Upd	t Database	Query	Time	Comment
	_	PGPB,USPT,EPAB,JPAB,DWPI(acetyl adj choline adj esterase	2005-04-	
			or choline adj esterase or	26	
		c	cholinomimetic) and	12:47:16	
1			substance or opioid or alcohol		
			or drug or nicotine or heroin)		
			adj3 (dependancy or		
			dependency or withdrawal or		
			abuse or overuse or inflence)		
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	•		or choline adj esterase) and	26	
1			substance or opioid or alcohol	12:45:00	
			or drug or nicotine or heroin)		
			adj3 (dependancy or		
* :			dependency or withdrawal or		- 17
2:0			abuse or overuse or inflence)		
<u>S10258</u>	<u>U</u>	PGPB,USPT,EPAB,JPAB,DWPI(
			or choline adj esterase) and	26	
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			nicotine or heroin) adj3		
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<u> </u>	<u>U</u>	PGPB,USPT,EPAB,JPAB,DWPI	or choline adj esterase) and	26	
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310230	<u>U</u>		esterase or choline adj esterase)		
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			drug or nicotine or heroin) adj3		8
			abuse		
S10255	U	PGPB,USPT,EPAB,JPAB,DWPIi		2005-04-	
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<u>\$10252</u>	<u>U</u>	PGPB,USPT,EPAB,JPAB,DWPI donepizil or aricept and	26 11:21:59 2005-04-
		(substance or opioid or drug or nicotine or heroin) adj3 abuse	26 11:17:25
<u>S10251</u>	<u>U</u>	PGPB,USPT,EPAB,JPAB,DWPI donepizil or aricept	2005-04- 26
S10250	U	PGPB,USPT,EPAB,JPAB,DWPI donepizil	11:16:55 2005-04-
			26 11:16:48
<u>S10249</u>	<u>U</u>	PGPB,USPT,EPAB,JPAB,DWPI aricept	2005-04- 26
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